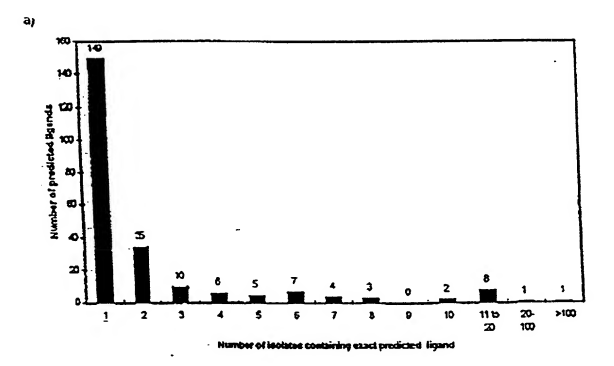
37.3



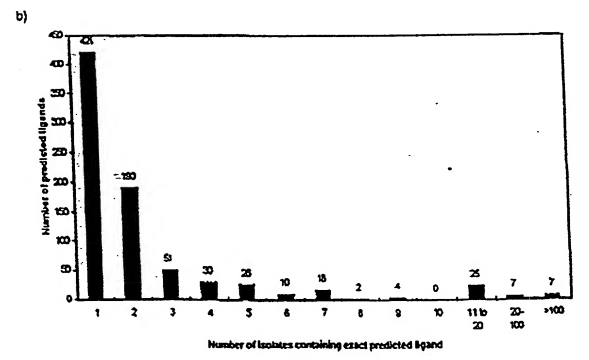


FIG. 2

Other SEQ ID NO.	-48487878787878787878787878
Other	××× × ××××××
Clade	× × × × ×
Clade	××× ××× ×××× × ×
Clade	× × ××× × ×
Clade B	×××××× ××××××× × × × ××××
Clade A	××× × ××××× × ×
Protein Number of Approximate Isolates Position in with Exact LAI AA sequence	gp120 - 120 gag - 215 gag - 215 gag - 216 RT - 354 nef - 170 nef - 170 gp120 - 452 gp120 - 420 gp120 - 420 gp120 - 420 gp120 - 420 gp120 - 420 gp120 - 215 gp120 - 215 gp120 - 215 gp120 - 215 gp120 - 215 gp120 - 275 gp120 - 275
Number of Isolates with Exact AA	159 159 171 171 183 171 171 183 183 183 183 183 183 183 18
Protein	Gag Gag Gag RT RT RT Gag Bry Bry Bry Bry Bry Bry Bry Bry Bry Bry
B27 Fold Increase (less than 1.3 not reported)	3.61 1.74 1.53 1.78 3.27
A2 Fold Increase (less than 1.3 not	1.33 1.33 1.63 1.54 ·
A2 EBP B27 EBP	0.00% 0.00% 0.00% 0.00% 0.00% 0.00% 0.00% 0.00% 0.00% 0.00% 14.22% 99.52% 89.62% 82.77% 82.11% 56.11% 56.11% 76.92% 56.14% 77.36% 67.49% 56.14% 76.92%
A2 EBP	55.68% 66.42% 66.42% 67.03% 77.52% 87.51% 90.92% 90.92% 90.00% 90.00% 90.00% 90.00% 90.00% 90.00% 90.00% 90.00% 90.00% 90.00% 90.00% 90.00% 90.00% 90.00%
Sequence	KLTPLCVTLN AEWDRVHPV SLENTVATL ELHPDKWTV GMDDPEREVL GMDDPEREVL GMDDPEREVL HLWRWSTMLL LLLTRDGGVN HLWRWSTMLL LLLTRDGGVN HLWRWSTMLL LLLTRDGGVN HLWRWSTMLL CREKQIN CREKQIN CREKQIN CREKQIN CREKQIN NSFEPPIHF RCSSNITGL VSFEPPIHF CREKQIVNM RSENITDN CREKQITY CREKQITY CREKQITY CREKQITY CRECGERIC RREAPODS RRESNITDN CREKQFIN KRUSIGPGR GCQCIEQL GREGGEREKY

FIG. 3

Project Outline

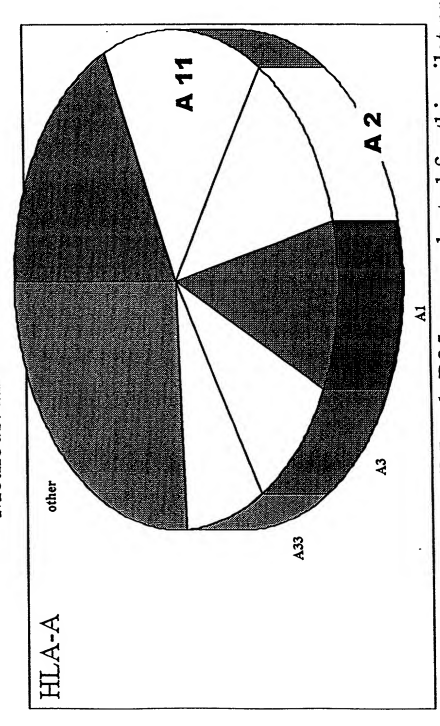
Indian HIV-1 sequences EpiMatrix evaluation and selection HLA allele selection

Binding assays (T2 stabilization)

CTL assays

Include CTL epitope in regional HIV-1 vaccine

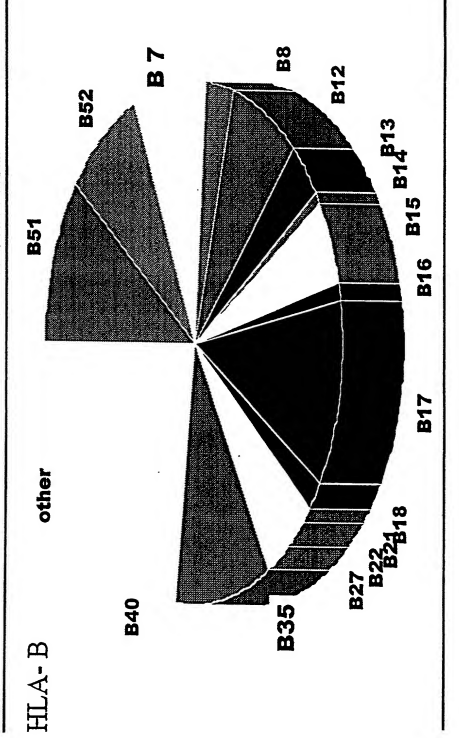
FIG. 4 Methods: HLA allele selection



HLA-A2 and -A11, -B7 and -B35 were selected for this pilot project due their prevalence of the HLA alleles in the Indian population....

FIG. 5

Methods: HLA allele selection



... and availability of cell lines for in vitro study.

FIG. 6

EpiMatrix Predictions and Binding Results: B 7 6 out of 7, and control peptide

Peptide seq. Used ge RPNNNTRKSI RPNNNTRKSI EN NPYNTPIFAL NPYNTPIFAL PC	gene strain ENV DID757 POL Solnd5	start-stop	L	•			
	Solnd5	The state of the s	% conserved	115	predicted EBP	AVG MFI (Zooug/mi)	strain start-stop % conserved GTL predicted EBH avg MFI (200ug/mi) avg told incr. (20ug/m
	Solnd5	183-192	75	Υ	8%	335.6	2.4
	DID747	61-70	90		20%	281.9	2.0
		481-490	80		17%	181.5	1,3
		375-384	20		18%	160.5	1.2
	DID747	182-191	7.		46%	248.5	1,8
	DID747	51-60	9		27%	373.8	2.7
	Solnd4	25-34	100		27%	314.7	2.3
		278-287	100		26%	378.4	2.7
~ - n & l	050	ENV DID747 T POL Soind4	Por	ENV DID747 T POL Soind4	ENV DID747 182-191 ENV DID747 51-60 T POL SoInd4 25-34	ENV DID747 182-191 ENV DID747 51-60 T POL SoInd4 25-34	ENV DID747 182-191 71 ENV DID747 51-60 100 T POL SoInd4 25-34 100 POL SoInd4 278-287 100

FIG. 7
EpiMatrix Predictions and Binding Results: B 35
7 out of 7 ... and control peptide

B37										
peptide #	septide	seq. Used	eueß	strain	start-stop	% conserved	CT	predicted EBP	avg MFI (200ug/ml)	start-stop % conserved CTL predicted EBP avg MFI (2004g/ml) avg fold indr. (204g/ml)
2	DVGDAYF	TVLDVGDAYF TVLDVGDAYF	집	Solnd4	114-123	100	>	4%	47.9	1,8
9	EPPFLWMGY	EPPFLWMGYE	S S	Solnd4	231-239	100		% 6	48.7	9 :
	VPVKLKPGM	VPVKLKPGMD	전	Solnd4	15-24	5		% 6	53.3	1.7
- α	CPKVTFDPI	CPKVTFDPIP		DID760	144-153	53		7%	35.0	<u>, i</u>
•	KPVVSTQLL	KPWSTOLL	ENV	DID747	182-191	7		%6	40.5	4.
	KPCVKLTPL	KPCVKLTPLC	ENS		51-80	9		11%	52.1	1.7
	GPKVKQWPL	GPKVKQWPLT	^o	Solnd4	25-34	5		11%	41.2	4
	YPGIKVRAL		PO.	POL Solnd4	278-287	100		7%	40.7	1.3

FIG. 8
EpiMatrix Predictions and Binding Results: A 2
3 out of 7 ... and control peptide

A2				-						
peptide #	peptide	seq. Used	gene	strain	start-stop	% conserved	CTL	start-stop % conserved CTL predicted EBP avg A	AFI (200ug/ml	avg fold incr. (20ug/ml
13	ILKEPVHGV	ILKEPVHGVY	POL L	Solnd4	316-325	80	>	%96	1604.2	1.6
14	QLPEKDSWTV	ALPEKDSWTV ALPEKDSWTV	POL	Solnd4	252-261	92		87%	1368.1	1.4
15	NEWTVYYGV NEWTVYYGV	NLWTVYYGV	ENS	GrD1024 32-41	32-41	67		84%	1716.9	8:
48	QMHEDVISI.	QMHEDVISLW	EN≤	DID747	37-46	9		78%	1413.1	1.4
4	KIEELREHLL	KIEELREHLL	PQ.	Solnd5	208-217	90		78%	889.9	6:0
18	DMVNQMHEDV	DMVNQMHEDV DMVNQMHEDV		DID747	33-42	2		77%	731.1	4.0
6	GLKKKKSVTV GLKKKKSVTV	GLKKKKSVTV	P _O	Solnd4	106-115	100		76%	1088.4	7
20	ELHPDKWTV	ELHPDKWTV ELHPDKWTVQ	P _Q	Solnd4	240-249	80		72%	1048.1	4.0

FIG 9

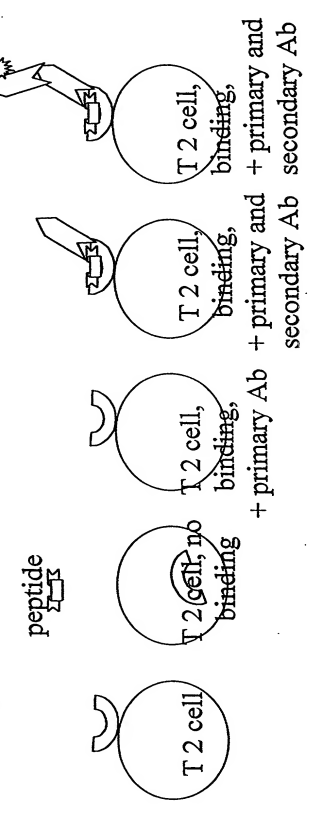
EpiMatrix Predictions and Binding Results: A 11 4 out of 7 ... and control peptide

A11										
peptide #	peptide	seq. Used	gene	strain	start-stop	% conserved	CTL	strain start-stop % conserved CTL predicted EBH avg	avg MFI (200ug/ml)	WFI (200ug/ml) avg fold incr. (20ug/ml,
21	IYQEPFKNLK	IYQEPFKNLK	POL	Solnd4	348-357	100	>	7%	677.5	3.1
22	VTFDPIPIHY	VTFDPIPIHY	ENS	DID780	147-158	53		22%	190.0	6.0
23	TVQCTHGIK	TVQCTHGIKP	ENS	DID747	174-183	59		44%	733.4	3.3
24	NTPIFALKKK	NTPIFALKKK	집	Solnds	64-73	09		44%	187.8	6.0
25	LVDFRELNK	LVDFRELNKR	P _Q	Solnd4	81-90	9		47%	755.2	3.4
70	PGMDGPKVK	PGMDGPKVKQ	Po	Solnd4	21-30	9		52%	193.8	0.7
27	GIPHPAGLKK	GIPHPAGLKK	PO L	Solnd4	100-109	100		62%	309.6	4.
28	FTTPDKKHQK	FTTPDKKHQK	전	Solnd4	221-330	18		63%	920.8	4.1

FIG. 10

Methods: T2 Binding Assay

TAP deficient cells. The stabilized MHC-peptide complex is detected Allele matched peptides stabilize MHC molecules on the surface of using Ab to the MHC and fluorescence labeled secondary Ab.



Clustering of putative MHC ligands in env

FIG. 11

